NIH -- W1 J0828FE

PAMELA GEHRON ROBEY

CSDB/NIDR/NIH Bldng 30 Rm 228 30 CONVENT DRIVE MSC 4320 BETHESDA, MD 20892

SUBMITTED: 2002-01-04 12:17:37 ATTN: 2002-01-08 11:11:07 PHONE: 301-496-4563 PRINTED:

REQUEST NO.: NIH-10102409
SENT VIA: LOAN DOC FAX: 301-402-0824 E-MAIL:

5433018

NIH Fiche to Paper Journal

TITLE: JOURNAL OF PEDIATRIC ORTHOPEDICS

Raven Press New York Ny PUBLISHER/PLACE:

VOLUME/ISSUE/PAGES: 1988 Sep-Oct;8(5):599-601 599-601

DATE: 1988

AUTHOR OF ARTICLE: Troop JK; Herring JA

TITLE OF ARTICLE: Monostotic fibrous dysplasia of the lumbar spine:

ISSN: 0271-6798

OTHER NOS/LETTERS: Library reports holding title, but not vol or yr

> 8109053 3049670 PubMed

SOURCE: W1 J0828FE CALL NUMBER:

REQUESTER INFO: AB424

DELIVERY: E-mail: probey@DIR.NIDCR.NIH.GOV

REPLY: Mail:

NOTICE: THIS MATERIAL MAY BE PROTECTED BY COPYRIGHT LAW (TITLE 17, U.S. CODE)

----National-Institutes-of-Health,-Bethesda,-MD-------

nours of

nington, 345. Parative

-bones. 3. Giant

cases, j csion of carative ant cell

A case 85;193;

ction of e granplasia.

W III

hands

outside

erature

. Hum

e bone bones,

carpal.

uloma

Surg

rative

iew of

Cam-

ma of

Intern

rative

I Clin

Case Report

Monostotic Fibrous Dysplasia of the Lumbar Spine: Case Report and Review of the Literature

J. Keith Troop, M.D., and John A. Herring, M.D.

Texas Scottish Rite Hospital and University of Texas Health Sciences Center, Dallas, Texas

summary: Fibrous dysplasia has been frequently reported to involve the spine in the polyostotic form, but only rarely has monostotic fibrous dysplasia been noted. in the only previously reported case involving the lumbar one, the disease was confined to the transverse process. The present case demonstrates monostotic fibrous dysplasia involving the vertebral body in addition to the posterior elements. The plain radiographic, computerized tomography, and histologic examinations are presented. **Key Words:** Lumbar spine—Monostotic fibrous dysplasia.

Fibrous dysplasia is a benign, relatively common fibroosseous lesion of bone. The earliest reports discussed only cases with involvement of multiple bones (1,10-12). Subsequently, three distinct forms of involvement were recognized (11,18) monostotic, polyostotic, and polyostotic with cutaneous and endocrine abnormalities (Albright syndrome) (1). The most frequent areas of involvement in each of these forms are ribs, proximal femur, tibia, skull, and maxilla (9,14,19,21). Spinal involvement is uncommon in the polyostotic form (10,11,14,20) and rare in the monostotic form (4,9,11,19,20).

CASE REPORT

The patient was referred for treatment of a lumbar lesion. She was a 12½-year-old white girl in good health until she fell from her bicycle. She sustained two phalangeal fractures and complained of midthoracic back pain. Radiographs of the spine demonstrated an expansile, lytic lesion of L3 (Fig. 1). A technetium bone scan showed increased uptake in the midlumbar region. Computerized tomography of the lumbar spine revealed a lytic, expansile lesion involving the right posterior body, pedicle, superior articular process, and right transferse process of L3 (Fig. 2). She was then referred for further evaluation.

On presentation to our institution 5 months after the initial trauma, the patient was asymptomatic. The medical history was significant for treatment as an infant for bilateral retinoblastoma with chemotherapy and local radiation followed by enucleation. At follow-up, she was considered to be disease free. Her general physical examination was within normal limits. She was premenarchal with no stigmata of endocrine abnormalities. The back was nontender and had a full range of motion. The

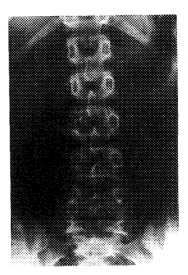


FIG. 1. Anteroposterior radiograph of the lumbar spine demonstrates an expansile, lytic lesion of L3 involving primarily the right posterior elements.

Address correspondence and reprint requests to Dr. J. A. Berring at Texas Scottish Rite Hospital, 2222 Welborn, Dallas, X 75219.

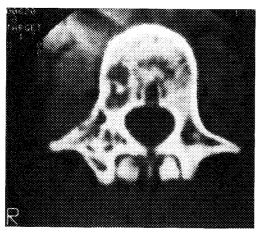


FIG. 2. Computerized tomography scan at the level of L3 demonstrates a lesion involving the right posterior body, right pedicle, right superior articular process, and right transverse process.

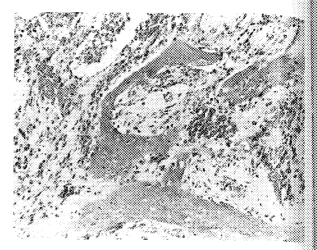
neurological examination was normal, and the initial laboratory examination was within normal limits. The presumptive diagnosis was aneurysmal bone cyst. Because of the history of malignancy as an infant and the atypical radiographic appearance of the lesion, a biopsy was performed.

Through a midline incision from L1 to the sacrum, the spine was exposed. The external appearance of the posterior elements was normal. The location was confirmed radiographically. The right superior facet and right superior portion of the transverse process of L3 were removed, and a cavity was entered. The cavity was surrounded by dense sclerotic bone with a thin membrane lining. There were sparse trabeculae transversing the cavity and no fluid. Following curettement of the lesion, autogenous bone graft obtained from the posterior iliac crests was packed in the cavity. A posterior fusion from L2 to L3 was performed. Microscopic examination showed woven bone with intervening fibrous tissue (Fig. 3). The pathological diagnosis was fibrous dysplasia.

The postoperative course was unremarkable. Before discharge, the patient was fitted with a posterior brace with anterior corset. The brace was removed at 11 weeks, at which time there was radiographic evidence of fusion (Fig. 4). When last seen 3 years after her initial injury, she was pain free with no scoliosis.

DISCUSSION

The earliest reports of fibrous dysplasia were primarily of severe polyostotic cases, most with the associated pigmentation and endocrine abnormalities constituting Albright syndrome (1,2-4,10-12,22). As more subtle forms of the bone pathology were appreciated, the monostotic form came to predominate in more recent series (9,14,19,20). Vertebral fibrous dysplasia is uncommon but not rare in polyostotic disease and in-



mber mic dis yem More mot mon we be series eriod. 1, 37 fright stated umbai spine angle ₹2-year

gadiog

sion o

firmed single

cothor

67 pat

had a

preser

pain r

the of

of C5

efter

the di

the s

which

of the

patho

was

const

invol

found

fibro

 Ω_{2}

and

Stuti

plasi

(n :

Hen

bral

222

clud

brot

the

this

por

inv

casi

teni

nos

the

SIO

Set

FIG. 3. Photomicrograph of the biopsy specimen reveals in regular trabeculae of woven bone with intervening fibrous tissue typical of fibrous dysplasia.

creases in frequency with increasing involvement of the rest of the skeleton. Therefore, the early cases actually have fairly frequent vertebral in volvement. Albright et al. (1) described five patients with severe polyostotic disease with the associated stigmata that now make up Albright syndrome. Three of these patients had vertebral involvement, one with L3 alone and the other two with multiple involvement. The patient of McCune and Bruch (12) had collapse of multiple vertebrae. The original article by Lichenstein (10) in 1938 stated that "lesions have noted roentgenographic cally on the vertebrae." One of the four polyostotic patients had vertebral involvement. Furst and Shapia (5), Albright and Reiferstern (2), Dockerty et al. (3), and Warrick (22) each reported on small

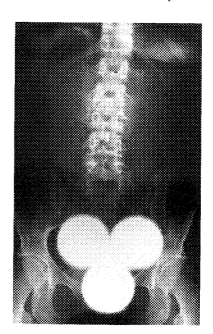


FIG. 4. Follow-up radiograph demonstrates fusion between L2 and L3. No scoliosis was apparent clinically.

mbers of patients, all with widespread polyosic disease and a high incidence of vertebral invenent.

More recent series have been predominated by monostotic form of the disease. Only two cases monostotic fibrous dysplasia above the sacrum we been reported (8,20). Harris et al. (8) reported eries of 90 patients who presented over a 30-year giod. Of the 50 cases that they described in de-37 were polyostotic (including those with Algight syndrome) and 13 were monostotic. They mated that of the polyostotic group, 14% had mbar spine involvement and 7% had cervical mine involvement. Of the monostotic group, a ngle case involved the spine. The patient was a year-old man who presented with low back pain. adiographs revealed an expanded radiolucent lesion of a transverse process of L4. Biopsy conrmed the diagnosis of fibrous dysplasia. The angle report of a monostotic lesion in the cerviothoracic spine is from Schlumberger (20). Of his patients with monostotic fibrous dysplasia, one ad a lesion of C4. The patient, a 20-year-old man, mesented with 11 months of local tenderness and pain radiating down the arms. Seven months before the onset of symptoms, he had sustained a fracture of C5 without neurologic sequelae. Four months after the fracture (three months before the onset of the described symptoms), he sustained a blow to the same area. He apparently had a lesion of C4, which was biopsied. No radiographs or descrition of the distribution in C4 is provided. The initial pathological diagnosis was giant cell tumor, which was revised to fibrous dysplasia on review by the consulting pathologist.

Several recent reports refer only briefly to spinal involvement. Reed (16) reviewed 25 patients and found 16 with monostotic and nine with polyostotic fibrous dysplasia. Vertebral fibrous dysplasia (L2-L5) occurred in only one polyostotic patient and none in the monostotic group. Firat and Stutzman (4) had one case of sacral fibrous dysplasia in each of his their main groups: monostotic (n = 15) and polyostotic (n = 9). The series of Henry (9) of 50 monostotic patients had no vertebral involvement. The report by Schajowicz (19) of 222 monostotic cases and 36 polyostotic cases included a sacral lesion in the monostotic group.

The present case demonstrates monostotic fibrous dysplasia of the lumbar spine. A review of the major series of fibrous dysplasia indicates that this lesion is unusual. The single previously reported case of monostotic lumbar fibrous dysplasia involved only the transverse process. The present case involved the posterior elements but also extended to the vertebral body. Although the diagnosis was suspected preoperatively, the rarity of the presentation necessitated biopsy. The progression of individual lesions in fibrous dysplasia slows

or halts at puberty (9,21). Malignant degeneration (13,17) and spinal cord compression (15) may occur; however, both have been reported only in severe polyostotic cases.

If the symptoms are minimal and the radiographic appearance is that of a benign lesion, surgical treatment is not mandatory. When there is diagnostic uncertainty, an excisional biopsy is indicated. Fusion is necessary only when removal of posterior elements has been extensive enough to produce instability.

REFERENCES

- Albright F, Butler AM, Hampton AO, Smith P. Syndrome characterized by osteitis fibrosa disseminata, areas of pigmentation and endocrine dysfunction, with precocious puberty in females. N Engl J Med 1937;216:727-46.
- Albright F, Reiferstern EC Jr. The parathyroid glands and metabolic bone disease: selected studies. Baltimore: Williams & Wilkins, 1948:393.
- Dockerty MB, Ghormley RK, Kennedy RLJ, Pugh DG. Albright's syndrome (polyostotic fibrous dysplasia with cutaneous pigmentation in both sexes and gonadal dysfunction in females). Arch Intern Med 1945;75:357-75.
- Firat D, Stutzman L. Fibrous dysplasia of the bone—a review of twenty-four cases. Am J Med 1968;44:421-9.
- Furst NJ, Shapia R. Polyostotic fibrous dysplasia: review of the literature with two additional cases. *Radiology* 1943; 40:501-15.
- Gaupp V. Pubertas praecox bei Osteodystrophia Fibrosa. Monatsschr Kinderheilkd 1932;53:312.
- Grabias SL, Campbell CJ. Fibrous dysplasia. Orthop Clin North Am 1977;8:771-83.
- 8. Harris WH, Dudley HR, Barry RL. The natural history of fibrous dysplasia. *J Bone Joint Surg* [Am] 1962;44:207-33.
- 9. Henry A. Monostotic fibrous dysplasia. J Bone Joint Surg [Br] 1969;51:300-6.
- 10. Lichtenstein L. Polyostotic fibrous dysplasia. Arch Surg 1938;36:874-98.
- 11. Lichtenstein L, Jaffe HL. Fibrous dysplasia of bone. Arch Pathol 1942;33:777-816.
- 12. McCune DJ, Bruch H. Osteodystrophia fibrosa. Am J Dis Child 1937;54:806-48.
- 13. Milgram JW. Malignant degeneration of polyostotic fibrous dysplasia of bone. Bull Hosp Joint Dis 1975;36:137.
- Mirra JM. Bone tumors, diagnosis and treatment. Philadelphia: Lippincott, 1980:130-3.
- Montoya G, Evarts C, Dohm D. Polyostotic fibrous dysplasia and spinal cord compression. A case report. J Neurosurg 1968;29:102-5.
- 16. Reed RJ. Fibrous dysplasia of bone. Arch Pathoi 1963;75:480-95.
- 17. Riddell DM. Malignant change in fibrous dysplasia. Report of a case. J Bone Joint Surg [Br] 1964;46:251-5.
- 18. Russell LW, Chandler FA. Fibrous dysplasia of bone. J Bone Joint Surg [Am] 1950;32:323-7.
- 19. Schajowicz F. Tumors and tumorlike lesions of bone and joints. New York: Springer-Verlag, 1981;478-90.
- Schlumberger HG. Fibrous dysplasia of single bones (monostotic fibrous dysplasia). The Military Surgeon 1946;99:504
 27
- Spint HJ, Dorfman HO, Fechner RE, Ackerman LV. Tumor of bone and cartilage. Washington, DC: Armed Forces Institute of Pathology, 1971:270-80.
- Warrick CK. Polyostotic fibrous dysplasia—Albright's syndrome. A review of the literature and report of four male cases, two which were associated with precocious puberty.
 J Bone Joint Surg [Br] 1949;31:175-83.

en reveals in ining fibrous

volvement the early rtebral in. d five paith the asbright synvertebral other two of McCune vertebrae. 0) in 1938 enographioolyostotic Furst and Dockerty d on small

on between